

FILE 'HOME' ENTERED AT 15:29:09 ON 07 NOV 2003

L1 744 (MEASLES OR MUMPS OR RUBELLA) (S) (CANCER OR TUMOR OR CARCINOMA)

(FILE 'HOME' ENTERED AT 15:29:09 ON 07 NOV 2003)

FILE 'MEDLINE, CAPLUS, BIOSIS, EMBASE' ENTERED AT 15:29:38 ON 07 NOV 2003

L1 744 S (MEASLES OR MUMPS OR RUBELLA) (S) (CANCER OR TUMOR OR CARCINO  
L2 29 S L1 AND ATTENUAT### (S) (MEASLES OR MUMPS OR RUBELLA)  
L3 416 S L1 AND (MUMPS OR RUBELLA)  
L4 156 S L3 AND RUBELLA  
L5 6 S L4 AND L2  
L6 5 DUP REM L5 (1 DUPLICATE REMOVED)  
L7 122 DUP REM L4 (34 DUPLICATES REMOVED)  
L8 88 S L7 NOT PY>1999  
L9 39 S L8 AND RUBELLA (S) (CANCER OR TUMORE OR CARCINOMA)  
L10 83 S L8 AND RUBELLA (S) (CANCER OR TUMOR OR CARCINOMA)  
L11 27 S L10 AND (MEASLES OR MUMPS)  
L12 0 S L2 AND (POINT-MUTATION OR POINT (A) MUTATION)  
L13 2 S L1 AND (POINT-MUTATION OR POINT (A) MUTATION)  
L14 7 S ATTENUAT#### (S) MEASLES AND (POINT-MUTATION OR POINT (A) MUT  
L15 2 DUP REM L14 (5 DUPLICATES REMOVED)  
L16 2 S L15 NOT L13

L6 ANSWER 1 OF 5 CAPLUS COPYRIGHT 2003 ACS on STN

AN 2002:240505 CAPLUS

DN 136:257218

TI A method for limiting the growth of **cancer** cells using an **attenuated measles** virus

IN Russell, Stephen James; Fielding, Adele; Peng, Kah-Whye; Grote, Deanna

PA Mayo Foundation for Medical Education and Research, USA

SO PCT Int. Appl., 76 pp.

CODEN: PIXXD2

DT Patent

LA English

FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	WO 2002023994	A1	20020328	WO 2001-US42259	20010921
	W:	AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PH, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM			
	RW:	GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG			
	AU 2001095063	A5	20020402	AU 2001-95063	20010921
PRAI	US 2000-668196	A2	20000922		
	WO 2001-US42259	W	20010921		

AB A method for treating **cancer** cells is provided comprising directly or systemically administering a therapeutically ED of an **attenuated measles** virus. In one embodiment, the therapeutically ED is from about 103 pfus to about 1012 pfus and is delivered by direct injection into a group of cancer cells or via i.v. injection.

RE.CNT 10 THERE ARE 10 CITED REFERENCES AVAILABLE FOR THIS RECORD  
ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 2 OF 5 EMBASE COPYRIGHT 2003 ELSEVIER INC. ALL RIGHTS RESERVED.  
on STN

AN 2002227841 EMBASE

TI Immunomodulatory vaccination in autoimmune disease.

AU Urbanek-Ruiz I.; Ruiz P.J.; Steinman L.; Fathman C.G.

CS Dr. C.G. Fathman, Division of Immunology, Center for Clinical Immunology, Stanford Univ. School of Medicine, 269 Campus Drive, Stanford, CA 94305, United States. cfathman@stanford.edu

SO Endocrinology and Metabolism Clinics of North America, (2002) 31/2 (441-456).

Refs: 84

ISSN: 0889-8529 CODEN: ECNAER

PUI S 0889-8529(01)00021-4

CY United States

DT Journal; General Review

FS 003 Endocrinology

026 Immunology, Serology and Transplantation

037 Drug Literature Index

LA English

SL English

AB The development of vaccines is arguably the most significant achievement in medicine to date. The practice of inoculation with the fluid from a sore to protect from a disease actually dates back to ancient China; however, with the introduction of Jenner's smallpox vaccine, and greater

understanding of the immune system, vaccines have become specific and systematic. Traditional vaccines have used killed pathogens (hepatitis A and the Salk polio vaccines), immunogenic subunits of a given pathogen (hepatitis B sub-unit vaccine), or live **attenuated** pathogens ( **measles, mumps, rubella**, Sabin polio vaccines) to generate protective immunity. Currently, a new generation of vaccines that use the genetic material of a pathogen to elicit protective immunity are being developed. Although the most widespread and successful use of vaccines today remains in the arena of infectious diseases, manipulations of immune responses to protect against **cancers**, neurologic diseases, and autoimmunity are being explored rigorously.

L6 ANSWER 3 OF 5 CAPLUS COPYRIGHT 2003 ACS on STN DUPLICATE 1

AN 2001:732865 CAPLUS

DN 136:277865

TI Tumor necrosis factor-.alpha./interleukin-10 balance in normal and cystic fibrosis children

AU Shmarina, Galina V.; Pukhalsky, Alexander L.; Kokarovtseva, Svetlana N.; Pukhalskaya, Daria A.; Shabalova, Lidia A.; Kapranov, Nikolai I.; Kashirskaja, Natalia J.

CS Laboratory of Immunogenetics, Research Centre for Medical Genetics, Moscow, 115478, Russia

SO Mediators of Inflammation (2001), 10(4), 191-197

CODEN: MNFLEF; ISSN: 0962-9351

PB Carfax Publishing

DT Journal

LA English

AB The balance between tumor necrosis factor-.alpha. (TNF-.alpha.) and interleukin-10 (IL-10) is important for immune homeostasis maintenance. Exuberant prodn. of TNF-.alpha. contributes to an overwhelming inflammatory response and tissue damage. Commonly, however, increase in TNF-.alpha. is counterbalanced by the simultaneous synthesis of an anti-inflammatory cytokine IL-10, which suppresses prodn. of many activating and regulatory mediators. Here, the relationships between TNF-.alpha. and IL-10 in the plasma of healthy school-children and cystic fibrosis (CF) patients have been investigated. Blood samples were obtained from 12 CF patients with chronic pulmonary disease and 18 healthy school-children vaccinated with live **attenuated rubella** vaccine. IL-10 and TNF-.alpha. were detd. in the plasma samples using com. available ELISA kits. Before vaccination, most healthy children (13 of 18) demonstrated superiority of pro-inflammatory TNF-.alpha. over anti-inflammatory IL-10 (TNF-.alpha./IL-10 >1). In these subjects, a pos. linear assocn. between the cytokine values was found. Vaccine challenge resulted in a marked redn. of the TNF-.alpha./IL-10 ratios. In addn., the correlation between the cytokine values disappeared. Such disturbance was related to a high rise of IL-10 levels after inoculation. On the contrary, in CF individuals, plasma cytokine values remained in strong linear assocn. independently of TNF-.alpha. or IL-10 predominance. No spikes were obsd. in the plasma levels of IL-10 in CF patients during a 6-mo observation period. There were no fundamental differences between CF and healthy children in the regulation of TNF-.alpha. and IL-10 secretion. Thus, immune quiescence seems to be assocd. with the predominance of TNF-.alpha., whereas immune disturbance is characterized by IL-10 superiority. The only abnormality that was found in CF patients consisted of their inability to produce unlimited IL-10 in response to antigenic stimuli.

RE.CNT 44 THERE ARE 44 CITED REFERENCES AVAILABLE FOR THIS RECORD  
ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 4 OF 5 EMBASE COPYRIGHT 2003 ELSEVIER INC. ALL RIGHTS RESERVED.  
on STN

AN 2001332616 EMBASE  
 TI [Vaccination].  
 IMPFUNGEN.

AU Graubner U.B.; Liese J.; Belohradsky B.H.  
 CS Dr. U.B. Graubner, K.klin./K.poklin. D. Haunerschen K., Klinikum Univ.  
 Munchen-Innenstadt, Abteilung Hamatologie und Onkologie, Lindwurmstr. 4,  
 80337 Munchen, Germany  
 SO Klinische Padiatrie, (2001) 213/SUPPL. 1 (A77-A83).  
 Refs: 38  
 ISSN: 0300-8630 CODEN: KLPDB2

CY Germany  
 DT Journal; Article  
 FS 016 Cancer  
 017 Public Health, Social Medicine and Epidemiology  
 026 Immunology, Serology and Transplantation  
 037 Drug Literature Index

LA German  
 SL English; German  
 AB Vaccination has been an important part of antiinfectious prophylaxis in  
 pediatric oncology comprising immunizations with special indication like  
 varicella vaccine and follow-up of routine immunizations after  
 chemotherapy and bone marrow transplantation (BMT). Studies from the last  
 decade demonstrate a loss of long term immunity to immunization  
 preventable disease in most patients with chemotherapy and BMT who had  
 received appropriate immunization before. So far routine vaccination  
 programs following intensive chemotherapy have not been studied  
 prospectively. Immunization programs following BMT have shown that  
 immunizations with tetanus toxoid, diphtheria toxoid, inactivated  
 poliovirus vaccine and influenza vaccine - given at least 12 months after  
 transplantation - are safe and effective. Vaccination with live  
**attenuated** trivalent vaccine against **measles**,  
**mumps** and **rubella** in patients without chronic "graft  
 versus host disease" (GVHD) and without ongoing immunosuppressive therapy,  
 performed 24 months after transplantation, proved to be safe too.  
 Recommendations have been published by 5 different official groups: (1.)  
 "Standige Impfkommision" (STIKO) and (2.) "Deutsche Gesellschaft fur  
 padiatrische Infektiologie" (DGPI) recommend varicella vaccine fur  
 children with leukemia in remission for at least 12 months, for children  
 with solid **tumors** and for patients getting an organ  
 transplantation. Both societies do not comment on the schedule of booster  
 vaccinations (with live **attenuated** vaccines) after the end of  
 chemotherapy and after BMT. (3.) "Qualitatssicherungsgruppe" der  
 "Gesellschaft fur padiatrische Onkologie und Hamatologie" (QS-GPOH)  
 recommends immunization with nonliving vaccines when the patient is off  
 therapy for at least 3 months and immunization with live  
**attenuated** vaccines when he is off therapy for at least 6 months.  
 This group does not comment on varicella vaccine which has been  
 controversial among pediatric oncologists. (4.) The "Infectious disease  
 working party of the European group for Blood and Marrow Transplantation"  
 (EBMT) recommends immunization with nonliving vaccines when the patient is  
 off transplantation for at least 12 months, without GVHD and without  
 immunosuppressive therapy. (5.) The "Guidelines for Preventing  
 Opportunistic Infections Among Hematopoietic Stem Cell Transplant (HSCT)  
 Recipients" published by the following american institutions and  
 societies: "Centers for Disease Control and Prevention", "Infectious  
 Diseases Society of America" and "American Society of Blood and Marrow  
 Transplantation" recommend that patients should be routinely revaccinated  
 after transplantation if they are off immunosuppressive therapy and do not  
 suffer from GVHD: beginning of vaccinations with nonliving vaccines in the  
 second year after HSCT, beginning of vaccinations with live  
**attenuated** vaccines in the third year after HSCT. Life-long

seasonal influenza vaccination is recommended for all HSCT candidates and recipients, beginning during the influenza season before HSCT and resuming >6 months after HSCT. IT would be appreciated if working groups of these societies could find consensus recommendations on open and controversial questions in the near future.

L6. ANSWER 5 OF 5 EMBASE COPYRIGHT 2003 ELSEVIER INC. ALL RIGHTS RESERVED.  
on STN  
AN 1999272066 EMBASE  
TI Viral Disease Update: Editorial.  
AU Severson J.L.; Tyring S.K.  
CS Dr. J.L. Severson, Department of Microbiology, University of Texas Medical  
Branch, Galveston, TX, United States  
SO Current Problems in Dermatology, (1999) 11/2 (41-70).  
Refs: 204  
ISSN: 1040-0486 CODEN: APDEBX  
CY United States  
DT Journal; General Review  
FS 013 Dermatology and Venereology  
037 Drug Literature Index  
LA English  
SL English  
AB Eight human herpesviruses have been identified and 10 antiviral drugs are  
Food and Drug Administration approved for their therapy. The herpesviruses  
are unique in that they all may cause primary infection, establish  
latency, and then reactivate if conditions of altered immunity develop.  
Herpes simplex virus type 1 is usually the cause of herpes labialis or  
cold sores. More than 85% of the population is seropositive for this  
virus, although only 20% to 40% of people have recurrent orolabial  
outbreaks. Herpes simplex virus type 2 (HSV-2) is the most common cause of  
genital herpes infection. One of every 5 people more than 12 years old is  
seropositive for HSV-2. The incidence has increased by 30% since 1976.  
Most people do not even know that they have been infected with the virus  
and that they can transmit the virus to a sexual partner. The antivirals  
acyclovir, valacyclovir, and famciclovir modify recurrent outbreaks of  
genital herpes and suppress outbreaks when taken on a daily basis. The  
side effects are comparable to those of placebo and no drug interactions  
have been identified. Valacyclovir may even prevent the transmission of  
HSV-2 from a seropositive person to a seronegative partner. Vaccines to  
modify recurrences of herpes simplex infections and prevent infections  
show promise. Varicella zoster virus causes varicella (chickenpox) and  
zoster (shingles). A live **attenuated** vaccine is available to  
prevent varicella. Acyclovir, valacyclovir, and famciclovir modify the  
very painful reactivation of varicella zoster virus in shingles.  
Famciclovir even decreases the length and severity of postherpetic  
neuralgia that can be debilitating in the elderly. Epstein-Barr virus, the  
fourth human herpesvirus, is associated with infectious mononucleosis and  
lymphoproliferative diseases in immunocompromised patients.  
Cytomegalovirus, the fifth human herpesvirus, is typically symptomatic  
only in neonates and the immunocompromised. Human herpesvirus type 6  
causes a mild, self-limited disease in childhood called sixth disease or  
exanthem subitum. No diseases have definitively been linked to the seventh  
human herpesvirus, but it is speculated that pityriasis rosea may be  
linked to this virus. The eighth human herpesvirus is associated with the  
violaceous lesions of Kaposi's sarcoma. The poxvirus manifests as  
molluscum contagiosum in children and sexually active adults and is most  
troublesome in the immunocompromised. Treatment with the antiviral  
cidofovir results in remarkable clearance of molluscum contagiosum in  
human immunodeficiency virus-infected individuals. Human papillomaviruses  
cause warts anywhere on the body. An estimated 30% to 50% of the sexually  
active population has genital human papillomavirus infection, called

condyloma acuminata. Several types of human papillomavirus are associated with anogenital **cancers**. Human papillomavirus DNA is found in greater than 93% of cervical **cancers**, the second most common cause of **cancer** death in women throughout the world. Earlier treatments of these infections involved nonspecific tissue destruction and significant recurrence rates. Podofilox gel and solution are safe and effective for self-treatment of genital warts. Treatment with imiquimod, an immune-response modifying agent that induces interferon alfa and other cytokines, results in clearance of warts and human papillomavirus DNA. There are no systemic side effects and local inflammatory reactions are tolerable for most patients. The viral exanthems, **measles** and **rubella**, have made a resurgence in recent years. Most cases in the United States have been linked to international importations. Hepatitis C is a parenterally transmitted RNA virus. Chronic hepatic disease occurs in 50% of patients with acute hepatitis C infection. Chronic disease can lead to cirrhosis and hepatocellular **carcinoma**. Dermatologic manifestations may be the only clinical evidence of underlying disease. Findings may include pruritus, porphyria cutanea tarda, vasculitis, salivary gland lesions, lichen planus, polyarteritis nodosa, urticaria, erythema nodosum, and erythema multiforme. A new combination therapy of ribavirin capsules and interferon alfa-2b recombinant for injection results in 45% of patients able to sustain reduced hepatitis C virus levels.

L11 ANSWER 7 OF 27 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC. on STN  
 AN 1998:113631 BIOSIS  
 DN PREV199800113631  
 TI Effect of immunosuppressive therapy on **measles**, **mumps**  
 and **rubella** (MMR) antibody in children successfully completing  
 treatment for **cancer**.  
 AU Feldman, S. [Reprint author]; Andrew, M.; Norris, M.; McIntyre, B.; Iyer,  
 R.  
 CS Univ. Miss. Med. Cent., Dep. Pediatrics, 2500 N. State St., Jackson, MS  
 39216-4505, USA  
 SO Abstracts of the Interscience Conference on Antimicrobial Agents and  
 Chemotherapy, (1997) Vol. 37, pp. 236. print.  
 Meeting Info.: 37th Interscience Conference on Antimicrobial Agents and  
 Chemotherapy. Toronto, Ontario, Canada. September 28-October 1, 1997.  
 ICAAC.  
 DT Conference; (Meeting)  
 Conference; Abstract; (Meeting Abstract)  
 Conference; (Meeting Slide)  
 LA English  
 ED Entered STN: 3 Mar 1998  
 Last Updated on STN: 3 Mar 1998

L11 ANSWER 13 OF 27 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC. on STN  
 AN 1981:96080 BIOSIS  
 DN PREV198121031076; BR21:31076  
 TI EPIDEMIOLOGIC REVIEWS VOL. 2.  
 AU SARTWELL P E; NATHANSON N  
 SO Epidemiologic Reviews, (1980) pp. VII+231P. SARTWELL, P. E. AND N.  
 NATHANSON (ED.). EPIDEMIOLOGIC REVIEWS, VOL. 2. VII+231P. JOHNS HOPKINS  
 UNIVERSITY PRESS: BALTIMORE, MD., USA; LONDON, ENGLAND. ILLUS.  
 Publisher: Series: Epidemiologic Reviews.  
 ISSN: 0193-936X. ISBN: 0-8018-2405-2(PAPER), 0-8018-2404-4(CLOTH).  
 DT Book  
 FS BR  
 LA ENGLISH  
 ED Entered STN: 28 Apr 1986  
 Last Updated on STN: 28 Apr 1986

L11 ANSWER 22 OF 27 EMBASE COPYRIGHT 2003 ELSEVIER INC. ALL RIGHTS RESERVED.  
 on STN  
 AN 91067916 EMBASE  
 DN 1991067916  
 TI Virus infections in children with cancer.  
 AU Morris D.J.  
 CS Department of Virology, Booth Hall Children's Hospital, Charlestown Road,  
 Manchester M9 2AA, United Kingdom  
 SO Reviews in Medical Microbiology, (1990) 1/1 (49-57).  
 ISSN: 0954-139X CODEN: RMEMER  
 CY United Kingdom  
 DT Journal; General Review  
 FS 007 Pediatrics and Pediatric Surgery  
 016 Cancer  
 047 Virology  
 037 Drug Literature Index  
 LA English  
 SL English  
 AB Recent improvements in outcome consequent on the use of intensive  
 chemoradiotherapy have emphasised the importance of viruses as causes of  
 morbidity and mortality in children with **cancer**. Varicella and  
**measles** are associated with high mortality rates in these children

because of the development of giant cell pneumonia or encephalitis. Hepatitis B and non-A non-B hepatitis may produce fatal chronic liver disease. Oral herpes simplex may progress to necrotic ulcers, secondary bacterial sepsis or haemorrhage. Rarely, herpes simplex disseminates to the liver, lungs, or brain. Cytomegalovirus, adenoviruses, parainfluenzaviruses, influenzaviruses, and respiratory syncytial virus occasionally produce severe disease in children receiving anti-**cancer** chemotherapy, but rotavirus, astrovirus, hepatitis A virus, **mumps rubella**, enterovirus, and rhinovirus infections are probably no more severe in these children than in normal children. Laboratory diagnosis of virus infections in children with malignant disease currently rests largely on virus isolation, electron microscopy, and detection of viral antigens by immunofluorescence. The availability of antiviral chemotherapy for herpesviruses, measlesvirus, and other respiratory viruses emphasises the need for rapid tests. Effective prophylaxis and therapy are available for herpes simplex, varicella and zoster. **Measles** remains a serious disease in children with **cancer**. Although prophylactic hyperimmune **measles** globulin and therapeutic ribavirin may control **measles** in these children in the future, improved uptake of **measles** vaccine in the general population is also necessary.

L11 ANSWER 25 OF 27 EMBASE COPYRIGHT 2003 ELSEVIER INC. ALL RIGHTS RESERVED.  
on STN

AN 83140190 EMBASE

DN 1983140190

TI Immunoblastic lymphadenopathy (IBL)-like T cell lymphoma.

AU Shimoyama M.; Tobinai K.; Minato K.; Watanabe S.

CS Dep. Clin. Lab., Natl. Cancer Cent. Hosp., Chuo-ku, Tokyo 104, Japan

SO Gann Monographs on Cancer Research, (1982) No. 28/- (121-134).

CODEN: GANMAX

CY Japan

DT Journal

FS 016 Cancer

025 Hematology

005 General Pathology and Pathological Anatomy

031 Arthritis and Rheumatism

026 Immunology, Serology and Transplantation

LA English

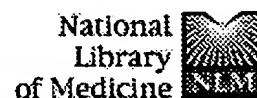
AB We proposed a new disease entity called 'Immunoblastic lymphadenopathy (IBL)-like T-cell lymphoma' in 1979. The present study represents an analysis of 14 cases. IBL-like T-cell lymphoma, although its clinical and morphological findings resemble immunodysplastic disease, IBL, angioimmunoblastic lymphadenopathy with dysproteinemia, lymphogranulomatosis x and polyclonal immunoblastosis, is a distinct peripheral T-cell lymphoma with suppressor/cytotoxic T-cell phenotype in adult; it is completely different from adult T-cell leukemia-lymphoma. The characteristics of this disease are summarized as follows: 1) the disease usually starts with generalized lymphadenopathy, frequently associated with high fever, skin rash and weakness; 2) lymphadenopathy is partially regressed by steroid hormone therapy, especially in the early phases of the disease; 3) frequent involvement of hepatosplenomegaly, but infrequent leukemic change and no thymic involvement; 4) poor prognosis; 5) marked male predominance; 6) polyclonal hypergammaglobulinemia; 7) Coombs test sometimes positive, occasionally associated with autoimmune hemolytic anemia and pure red cell aplasia; 8) elevation of various anti-virus titer (**measles**, **rubella**, varicella, and Epstein-Barr virus (EBV), and/or of anti-toxoplasma titer; 9) leucocytosis with neutrophilia, lymphocytopenia and atypical plasmacytoid cells; 10) no endemic distribution of the patients' birthplace; 11) multifocal or diffuse neoplastic proliferation of immunoblasts, large lymphoid cells and/or



so-called 'pale cells' with angioimmunoblastic and granulomatous lesions, zonal proliferation of plasma cells, disappearance of germinal center, deposition of amorphous acidophilic interstitial material and depletion of small lymphocyte; the patient is often diagnosed as IBL or angioimmunoblastic lymphadenopathy (AILD) at initial biopsy, so serial examinations must be indicated; 12) neoplastic cells express the T-cell nature of suppressor/cytotoxic T-cell phenotype; 13) surface and cytoplasmic immunoglobulins of lymph node cells are not monoclonal, but polyclonal. It is necessary to disclose why IBL-like T-cell lymphoma is frequently associated with variegated and brilliant clinical manifestations such as fever, skin rash, polyclonal hypergammaglobulinemia, elevation of various anti-virus titer and autoimmune mechanism, and granulomatous lesions in lymph nodes, despite the **tumor** cells having suppressor/cytotoxic T-cell phenotype.

L11 ANSWER 17 OF 27 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC. on STN.  
 AN 1977:183154 BIOSIS  
 DN PREV197764005518; BA64:5518  
 TI THE LIFETIME HEALTH MONITORING PROGRAM A PRACTICAL APPROACH TO PREVENTIVE MEDICINE.  
 AU BRESLOW L; SOMERS A R  
 SO New England Journal of Medicine, (1977) Vol. 296, No. 11, pp. 601-608.  
 CODEN: NEJMAG. ISSN: 0028-4793.  
 DT Article  
 FS BA  
 LA Unavailable  
 AB Current patterns of health care and its financing need to be improved by the incorporation of cost-effective and health-effective preventive measures. As a stimulus for further development, a Lifetime Health-Monitoring Program is proposed that uses clinical and epidemiologic criteria to identify specific health goals and professional services appropriate for 10 different age groups. During infancy, e.g., 7 immunizations [diphtheria, tetanus, pertussis, **measles**, **mumps rubella**, polio]; tests to detect anemia, hemorrhagic diseases, phenylketonuria and developmental deficiencies; and routine prophylaxis and gonorrheal ophthalmia are recommended. In the age group 40-59, tests for hypertension and cervical, mammary and gastrointestinal cancer, and control obesity and smoking are in order. The cost of such preventive measures, which should not be prohibitive, must be covered by health-insurance programs, whether based on fee-for-service or capitation. This program, by incorporating prevention into day-to-day care, should strengthen the patient-physician relation.

L16 ANSWER 2 OF 2 MEDLINE on STN  
 AN 96177164 MEDLINE  
 DN 96177164 PubMed ID: 8599233  
 TI The nonstructural C protein is not essential for multiplication of Edmonston B strain measles virus in cultured cells.  
 AU Radecke F; Billeter M A  
 CS Institut fur Molekularbiologie, Abteilung 1, Universitat Zurich, Honggerberg, Switzerland.  
 NC 5 R01 AI35136 (NIAID)  
 SO VIROLOGY, (1996 Mar 1) 217 (1) 418-21.  
 Journal code: 0110674. ISSN: 0042-6822.  
 CY United States  
 DT Journal; Article; (JOURNAL ARTICLE)  
 LA English  
 FS Priority Journals  
 OS GENBANK-Z66517  
 EM 199604  
 ED Entered STN: 19960506  
 Last Updated on STN: 19960506  
 Entered Medline: 19960425  
 AB Measles virus (MV) is a highly contagious agent which causes a major health problem in developing countries. Efficacious and safe live **attenuated** vaccine strains are available, but for the elimination of **measles** a better knowledge about the molecular biology of MV appears crucial. Whereas the roles of the six structural proteins in the replication cycle are known, the functions of the two nonstructural proteins C and V are unclear, which is also true for related viruses. In vitro studies implicating Sendai virus suggest that the C protein might be involved in downregulating viral mRNA synthesis (J. Curran, J.B. Marq, and D. Kolakofsky, Virology 189, 647-656, 1992). However, not all members of the Paramyxovirinae subfamily encode this protein, raising the question about its importance for the viral replication cycle. Taking advantage of a recently developed reverse genetics system allowing MV recovery from cloned DNA (F. Radecke, P. Spielhofer, H. Schneider, K. Kaelin, M. Huber, C. Dotsch, G. Christiansen, and M.A. Billeter, EMBO J. 14, 5773-5784, 1995), the question was addressed whether the C protein is essential for the life cycle of MV. A plasmid was constructed to produce a derivative of the Edmonston B vaccine strain, MV C- EdB, having its C reading frame silenced by two **point mutations**. The C- mutant MV could indeed be rescued, and it multiplies in cultured cells without obvious impairment.



Entrez	PubMed	Nucleotide	Protein	Genome	Structure	PMC	Journals	
Search	PubMed	for rubella AND vaccine AND (cancer or tumor)					Preview	Go
		✓ Limits	Preview/Index	History		Clipboard	Details	

- Search History will be lost after eight hours of inactivity.
- To combine searches use # before search number, e.g.; #2 AND #6.
- Search numbers may not be continuous; all searches are represented.

Entrez PubMed

Search	Most Recent Queries	Time	Result
#15	Search rubella AND vaccine AND (cancer or tumor) Field: Title/Abstract, Limits: Publication Date to 2000/09/22	15:20:15	<u>7</u>
#14	Search MMR AND vaccine AND (cancer or tumor) Field: Title/Abstract, Limits: Publication Date to 2000/09/22	15:20:07	<u>0</u>
#13	Search MMR AND (cancer or tumor) Field: Title/Abstract, Limits: Publication Date to 2000/09/22	15:19:43	<u>171</u>
#12	Search MMR same cancer Field: Title/Abstract, Limits: Publication Date to 2000/09/22	15:19:31	<u>8</u>
#11	Search Measles AND point-mutation Field: Title/Abstract, Limits: Publication Date to 2000/09/22	14:28:02	<u>2</u>
#9	Search Measles AND point mutation Field: Title/Abstract, Limits: Publication Date to 2000/09/22	14:27:40	<u>2</u>
#8	Search Measles AND attenuate AND point mutation Field: Title/Abstract, Limits: Publication Date to 2000/09/22	14:27:33	<u>0</u>
#4	Search measles AND F AND H Field: Title/Abstract, Limits: Publication Date to 2000/09/22	10:05:45	<u>99</u>
#1	Search measles AND F AND H Field: Title, Limits: Publication Date to 2000/09/22	09:53:44	<u>2</u>

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